

## **EU Declaration of Conformity**

We hereby declare under our sole responsibility that the Primus system meets the relevant provisions of the following European Union Directives:

- Council Directive 93/42/EEC of 14 June 1993 concerning medical devices as amended by Directive 2007/47/EC (MDD)
- Directive 2006/42/EC of the European Parliament and of the Council of 17 May 2006 on machinery as amended by Regulation (EU) 2019/1243
- Directive 2011/65/EU of the European Parliament and of the Council of 8 June 2011 on the restriction of the use of certain hazardous substances in electrical and electronic equipment (RoHS)

The Primus has undergone a conformity assessment procedure required by the MDD and is manufactured in harmony with the Technical Documentation compiled as defined in the relevant Directives and retained by BTE.

Product information in regard to the MDD and RoHS Directives:

Manufacturer	BTE Technologies 7455-L New Ridge Road Hanover, MD 21076, USA www.btetechnologies.com	Telephone: 410.850.0333 Email: Service@btetechnologies.com.
Product Identification	Device Trade Name: PrimusRS Device Name: Primus	S Model: PrimusRS (PRRS)
UDI-DI	10850390007243	
EMDN (CND) code	Z120616 - PHYSICAL THERAPY AND REHABILITATION SYSTEMS	
Intended Purpose	The Primus is intended to be used for musculoskeletal strength testing and exercise. Applications include physical rehabilitation and sports therapy. The system is intended to measure strength, increase muscle strength and endurance, and track patient progress through the process. It may be used for upper extremity, lower extremity, and trunk muscle weakness.	
Device Classification (MDD)	Class I	
Classification Rule (MDD)	Rule 12	



Route to Compliance (MDD)	Annex VII of the Medical Devices Directive	
Device Classification (MDR)	Class IIa	
Classification Rule (MDR)	Rule 11	
CE Marking Provision	Under Medical Device Regulation (EU) 2017/745 (MDR), the device will be up-classified to class IIa due to changed software classification rules. Based on the MDR Article 120 §3, the PRIMUS can be placed on the EU market as a class I device until May 26, 2024 provided that the device	
	<ul> <li>will continue to comply with the MDD,</li> </ul>	
	<ul> <li>there will be no significant changes in the design and intended purpose, and</li> </ul>	
	<ul> <li>the device will comply with the MDR requirements for post market surveillance, vigilance, and registration of economic operators and of devices</li> </ul>	
Authorized Representative  EC REP	Emergo Europe  Prinsessegracht 20  2514 AP, The Hague The Netherlands  Telephone: +31.70.345.8570  Emails: EmergoEurope@ul.com  EmergoVigilance@ul.com	

The device is CE marked since 2004.

Signed for on behalf of BTE Technologies

Ewa Kaczanowska

PRRC/Regulatory Manager

Wacnanowska

BTE Technologies

Hanover, MD

May 20, 2021